Topics in Molecular and Structural Biology

OLIGODEOXYNUCLEOTIDES

Antisense Inhibitors of Gene Expression

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Control of Gene Expression by Oligodeoxynucleotides Covalently Linked to Intercalating Agents and Nucleic Acid-cleaving Reagents

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Introduction

information processing: transcription of DNA into premessenger RNAs; splicing of mRNA precursors; post-transcriptional modifications of mRNAs Gene expression in all living organisms is controlled at different steps of (capping, polyadenylation); transfer of mRNAs from the nucleus to the cytoplasm; translation of mRNA; mRNA stability In most cases this regulation is achieved by proteins that bind to specific regions of DNA or RNA and either block or stimulate the enzymatic processes (see Hélène and Lancelot, 1982, for a review). Recently it has been shown that small RNAs could play a role similar to that of regulatory proteins. Upon hybridization with a messenger RNA, these regulatory RNAs may alter the translation process or induce premature termination of transcription (see Green et al., 1986, for a review). These regulatory processes have been originally observed in bacteria (Green et al., 1986) but they might also occur in eukaryotes (Heywood, 1986). The discovery of regulatory RNAs has been the starting point for the design of 'antisense' RNAs. By inserting a gene fragment close to a strong promoter in the reverse orientation as compared with that of the gene itself, the non-template strand of the gene fragment is mRNA translation or other post-transcriptional processes such as splicing now used as a template by RNA polymerase. As a consequence this 'antisense' transcript is fully complementary to the mRNA. This might block or mRNA migration from the nucleus to the cytoplasm (Kim and Wold,

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sequences to alter gene expression was put forward in several laboratories (Paterson et al., 1977; Stephenson and Zamecnik, 1978; Summerton, 1979; Jayaraman et al., 1981; Trudel et al., 1981; Asseline et al., 1983, 1984a, b; for reviews see also Knorre and Vlassov, 1985; Hélène et al., 1985; Hélène, 1987; Stein and Cohen, 1988; Toulmé and Hélène, 1988). It has been shown that oligodeoxynucleotides complementary to mRNAs could block transation in acellular systems, in microinjected Xenopus oocytes and in cells in culture. There is an obvious need for developing new families of gene reguatory substances that could be used in vivo to control the expression of undesirable genes, such as oncogenes, or to inhibit the development of viruses or parasites. The application of oligodeoxynucleotides to in vivo studies faces two main problems: (1) their penetration into living cells in The idea of using synthetic oligonucleotides complementary to RNA culture is limited; (2) their sensitivity to nucleases makes their lifetime very short (Cazenave et al., 1987b).

Several attempts have been made to overcome these two difficulties. The phosphodiester backbone of the oligodeoxynucleotide can be changed to a methylphosphonate backbone; the loss of negative charges makes these oligophosphonates more efficient in penetrating through the cell membranes and much more resistant to nucleases (Miller et al., 1983). The phosphate group can be replaced by a phosphorothioate (Marcus-Sekura et al., 1987); these oligophosphorothioates are much more resistant to nucleases than natural oligonucleotides. Attachment of oligonucleotides to polymers such as poly-t-lysine increases the efficiency of penetration and makes oligonucleotides active at much lower concentrations (Lemaître et al., 1987).

new families of specific gene regulatory substances. A nucleic acid base tide with its target sequence can be increased by covalent attachment of an 1984a,b). In addition, the intercalating agent endows the oligonucleotide which can be activated to modify the target sequence by either chemical or cal modification of the bases at the binding site of the oligonucleotide should prevent translation of the mRNA. In addition, the oligonucleotide can be modified in such a way as to make it more resistant to nucleases, e.g. This review summarizes the approach we have been following to design sequence can be easily recognized by an oligonucleotide of complementary with a higher penetration across cell membranes and stabilizes it against 3'-1987). The other end of the oligonucleotide can be substituted by a reagent Praseuth et al., 1987, 1988a). Specific cleavage of a mRNA target or chemiby substituting synthetic α -anomers of nucleotides for the natural β sequence. The stability of the mini-double helix formed by an oligonucleophotochemical activation (Boidot-Forget et al., 1986; Le Doan et al., 1987a; intercalating agent at one end of the oligonucleotide (Asseline et al., 1983, or S'exonucleases, depending on the attachment site (Verspieren et al.,

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DNA by binding to the major groove. Therefore oligonucleotides can be Oligodeoxynucleotides can recognize not only mRNAs but also duplex

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used to control gene expression at the transcriptional level. Oligonucleotides carrying a reactive group can induce irreversible reactions in duplex DNA, including double-strand cleavage.

2 Oligodeoxynucleotides as Anti-messengers

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Specificity of Oligonucleotide Targeting to Unique Sequences

Targeting to Genomic DNA

nize a single specific sequence in a genome can be calculated on the basis of The minimum size that an oligonucleotide should have in order to recogdifferent assumptions.

ized by a fraction f of A.T base pairs (f = [A.T]/[A.T] + [G.C]), the probability (p_0) of finding a sequence of n nucleotides is given by Equation Assuming a statistical distribution of base pairs in a genome charac

$$p_0 = [(f/2)]^{(a+t)} \times [(1-f)/2]^{(g+c)}$$
 (1)

sines in the oligonucleotide (n = a + t + g + c). The number (Q) of identical where a, t, g, c are the numbers of adenines, thymines, guanines and cytosequences of n nucleotides in a genome containing N base pairs is given by Equation (2):

$$Q = p_0 \times 2N \tag{2}$$

For E. coli, which contains about 4.5×10^6 base pairs in its genome with where the factor 2 accounts for the presence of the two strands in DNA. equal numbers of A.T and G.C base pairs (f = 0.5), the minimal length (n)that an oligonucleotide should have in order to find a single complementary sequence $(Q \le 1)$ is n = 12. In human cells, with $N = 4 \times 10^9$ and cleotide contains only Gs and Cs to n = 19 if the oligonucleotide contains $f \approx 0.6$, the minimal length is calculated to vary from n = 15 if the oligor only As and Ts.

The above calculation assumes a statistical distribution of base pairs in the genome, which, of course, is not correct. Analyses of nearest-neighbour frequencies have shown that the dinucleotide CpG is underrepresented in eukaryotic genomes. The probability of finding an oligonucleotide sequence in a genome can be calculated on the basis of nearest-neighbour frequencies using a first-order Markov chain. According to Markov's theory, this probability is equal to the product of probabilities of all overlapping dinucleotides in the sequence divided by the product of probabilities of shared mononucleotides. For example, the probability of finding the sequence CATCGT is given by Equation (3):

$$p_1(\text{CATCGT}) = \frac{p(\text{CA}) \times p(\text{AT}) \times p(\text{TC}) \times p(\text{CG}) \times p(\text{GT})}{p(\text{A}) \times p(\text{T}) \times p(\text{C}) \times p(\text{G})}$$
(3)

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where p(XY) is the probability of finding the sequence XpY, and p(X) is the probability of finding nucleotide X.

A second-order Markov chain can be used if the probability of finding trinucleotides (p(XYZ)) is known, as shown in Equation (4):

$$p_2(\text{CATCGT}) = \frac{p(\text{CAT}) \times p(\text{ATC}) \times p(\text{TCG}) \times p(\text{CGT})}{p(\text{AT}) \times p(\text{TC}) \times p(\text{CG})}$$
(4)

Higher-order Markov chains can be used if the frequency of longer sequences is known (tetranucleotides, pentanucleotides . . .). With the accumulation of sequence data, such calculations should become more and more accurate in predicting the probability of finding an oligonucleotide sequence in unknown regions of the genome.

ested in using oligonucleotides to specifically regulate gene expression, the tide but rather that of finding the complementary sequence. These two probabilities are obviously equal if the genome is the target, owing to the complementarity of the two strands in the DNA double helix. This is no longer true if the oligonucleotide is targeted to a messenger RNA. However, the only dinucleotide that significantly appears less frequently than calcu-Table 7.1 gives the probability of finding the decanucleotide sequence GGCATCGTCG in the E. coli and human genomes, according to equations (1) and (3). This sequence is found in the murine and human c-myc genes and was used in our laboratory to inhibit myc mRNA in vitro translation (see below). In E. coli the calculated probability does not markedly change with the mode of calculation. Obviously this is not the case for the human genome, because the chosen sequence contains two CpG dinucleotides which are underrepresented as compared with a statistical distribution. Equation (3) should be preferred to calculate the probability of finding any oligonucleotide sequence in eukaryotic genomes. Since we are interprobability that should be calculated is not that of finding the oligonucleolated on a statistical basis is CpG in eukaryotes. Its complementary sequence is also CpG. Therefore, the calculated probabilities will not be very different if an oligonucleotide or its complementary sequence is con-

Calculated probability of finding the sequence GGCATCGTCG in E. coli and human genomes calculated on the basis of zero-order and first-order Markov chains (equations 1 and 3, respectively) Table 7.1

Human	4.1×10^{-7} 0.22×10^{-7}
E. coli	9.5×10^{-7} 11×10^{-7}
	Po 9

Equation (1).

The frequency of nucleotides and dinucleotides was obtained from the Handbook of Biochemistry and Molecular Biology.

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Targeting to Messenger RNAs

asually transcribed, even though there is increasing evidence that divergent Only a small fraction of the eukaryotic genome is transcribed into messenger RNA in a living cell at a given time. Only one strand of the DNA is transcription of both DNA strands takes place in eukaryotes. It has been lower in the mRNA population than in DNA, since the target size drops from 4×10^9 to 2×10^7 units in human cells. On this assumption the minimal length that an oligonucleotide should have in order to find a single arget at the mRNA level is reduced: from 15 to 11 if the oligonucleotide ing several CpG dinucleotides might be chosen quite short without losing more than once, on a statistical basis, in the human mRNA population (the calculated probability is 0.22×10^{-7} and the target mRNA complexity estimated that about 0.5% of the genomic DNA is transcribed into mRNA. Therefore, the probability of finding an oligonucleotide sequence is much show that the decanucleotide sequence GGCATCGTCG should not occur (see above), the length of an oligonucleotide targeted to a sequence containthe specificity of mRNA recognition. The results presented in Table 7.1 contains only Gs and Cs and from 19 to 15 if it contains only As and Ts Owing to the low frequency of CpG dinucleotides in the human genol amounts to 2×10^7 nucleotides; see above).

When a viral RNA or a viral mRNA is chosen as a target for oligonucleotides, it should be kept in mind that codon usage is usually different from that of the host cell. There might be a difference in trinucleotide and tetranucleotide frequencies between the virus and its host. This difference can be taken into account when choosing a target sequence.

The conclusion that can be drawn from the above considerations is that a high specificity of oligonucleotide binding to mRNAs can be achieved with quite short oligonucleotides. Using short oligonucleotides provides additional advantages.

- (1) The probability of forming intramolecular secondary structures (hairpins) is lower in short oligonucleotides.
- (2) The probability of finding the target sequence in an accessible region of that RNAs adopt folded conformations. Complementary sequences which can be far apart in the primary sequence may form duplex structures. Hairpins may fold on themselves to engage in tertiary interactions a messenger RNA or a viral RNA is increased. It should be remembered (as observed, e.g., in the folded clover-leaf structure of tRNAs).

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- branes is expected to decrease when the length of the oligonucleotide increases. Therefore, short oligonucleotides could penetrate better inside The penetration of negatively charged oligonucleotides across cell memliving cells. 3
 - If oligonucleotides have to be chemically modified in order to make them, e.g., more resistant to nucleases, chemical synthesis and purifi-4

^b Equation (3).

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cost less!). These points might be of special interest if practical cation should be easier to achieve for short molecules (also they should (therapeutical) applications are contemplated.

The specificity of interaction will be higher with shorter oligonucleotides Studies on inhibition of gene expression in living cells are carried out under well-defined conditions of temperature, ionic concentrations, etc., 37 °C. The important parameters which should be considered in closely related sequences at this temperature (which determines the specificity of inhibition). The free energy cost due to a mismatch does when the mismatch is located close to the oligonucleotide end which is provided that the target (complementary) sequence is found only once. which are imposed upon the experimentalist by the species under investgation. For example, experiments on human cells are performed at choosing oligonucleotide length are the stability (the free energy) of binding to the target sequence at 37 °C and the discrimination between not change appreciably when an oligonucleotide is elongated, except elongated. But its relative contribution to complex destabilization of course decreases when the oligonucleotide length increases. (5)

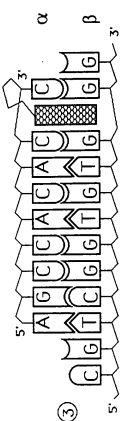
An important component in the biological activity of oligodeoxynucleotides is RNase-H, an enzyme which cuts RNA within the region hybridized to the oligodeoxynucleotide (see below). This enzyme recognizes duplexes as short as 4 base pairs (Donis-Keller, 1979) and could be complementarity. Such an effect should be minimized by using short responsible for non-specific inhibition of gene expression due to partial oligonucleotides. 9

Oligonucleotides Covalently Linked to Intercalating Agent

Increased Binding to Target Sequences

The above calculations suggest that short synthetic oligonucleotides can be designed to bind to a single nucleic acid target. However, a short oligonucleotide might not have a strong enough affinity towards its target sequence if the number of base pairs involved is too small. There are different ways of increasing this affinity. We have chosen to covalently link an intercalating two consecutive base pairs of double-stranded DNA. They bind much more occur in the mini-double helix formed when the oligonucleotide is bound to cyclic aromatic molecules that insert their planar aromatic ring between tide and the intercalating agent is appropriately chosen, intercalation can agent to one (or both) oligonucleotide end(s). Intercalating agents are polyits complementary sequence (Figure 7.1). To a first approximation the free Bridge-Intercalator) should be the sum of the free energy for binding the weakly to single-stranded structures. If the linker between the oligonucleoenergy of binding of the composite molecule (ONBI, for OligoNucleotideoligonucleotide to its complementary sequence (ΔG_{ON}) and that for inter-

'n <u>G</u> 9 0 6 6 () 9 6 9 <u>G</u> 9 'n വ



'n

വ

positions of the intercalating agent. In 3 an oligonucleotide with synthetic a-anomers is shown Figure 7.1 Schematic representation of an oligonucleotide covalently linked to an inter-calating agent via its 3' end (rectangle) when bound to a complementary sequence. In 1 ay an oligonucleotide with natural eta-anomers of the nucleotides is depicted with two diffe bound in a parallel orientation with respect to its complementary sequence

calation (ΔG_1), corrected for an entropy term ($T\Delta S_m$) taking into account the restricted configurational space available to the intercalating agent when it is covalently linked to the oligonucleotide:

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$$\Delta G_{\text{ONBI}} = \Delta G_{\text{ON}} + \Delta G_{\text{I}} - T\Delta S_{\text{m}} \tag{5}$$

Since ΔS_m in Equation (5) is positive, the association constant for the ONBI $(K_{ONBI} = \exp - \Delta G_{ONBI}/RT)$ should be at least the product of the association constants for the oligonucleotide and the intercalating agent.

$$K_{\text{ONBI}} = \alpha K_{\text{ON}} \times K_{\text{I}} \tag{6}$$

with $\alpha = \exp(\Delta S_m/R) > 1$.